Ellipse[™] VR

Single-Chamber Implantable Cardioverter Defibrillator (ICD)

Product Highlights

- MRI-ready device will allow patients to safely undergo an MRI scan when used in combination with an MR Conditional lead^{1,2}
- Improved shape with reduced volume and thickness
- Parylene coating for improved abrasion resistance
- DynamicTx[™] Over-Current Detection Algorithm automatically changes shock configurations to ensure delivery of high voltage therapy when high current is detected
- Cold Can programmability provides an additional RV-SVC Shock Configuration to decouple the can from the shocking vector parameters in cases of lead problems
- ShockGuard[™] technology with DecisionTx[™] programming designed to reduce inappropriate therapy and minimise the need for programming adjustments at implant
- Low Frequency Attenuation filter designed to enhance sensing performance and may reduce the possibility of oversensing T-waves
- SenseAbility[™] feature provides flexibility to fine-tune programming around T-wave oversensing without decreasing sensitivity
- DF4 connector designed to streamline defibrillation connections into a single terminal pin and reduce the number of set screws
- CorVue[™] congestion monitoring feature monitors the intrathoracic impedance in multiple vectors for improved accuracy, and it provides the option for both patient and physician alerts
- SecureSense[™] RV lead noise discrimination detects sustained and short bursts of lead noise that would otherwise go unnoticed or potentially lead to one or more inappropriate shocks
- Far Field MD[™] morphology discrimination improves SVT and VT discrimination for reduced inappropriate therapies
- Antitachycardia pacing (ATP) while charging and prior to charging in the VF zone further extends the programming options for terminating tachyarrhythmias without a high-voltage shock
- ST monitoring capability provides unprecedented, continuous insight into significant ST shift events and associated ventricular arrhythmias through enhanced monitoring of iEGM and ST-segment as a diagnostic tool to help guide appropriate clinical action
- 36 J delivered energy safety shock option can provide a greater DFT safety margin
- DeFT Response[™] technology offers the most noninvasive options for managing high DFTs
- QHR^{™†} chemistry battery provides greater capacity for enhanced longevity and improved charge time performance compared to previous SVO batteries

Ordering Information

Contents: Cardiac pulse generator

Model Number	Dimensions (H x W x T, mm)	Weight (g)	Volume (cc)	Connector Defibrillation	Connector Sense/Pace
CD1377-36C	68 x 51 x 12	66	31	DF1	IS-1
CD1377-36QC*	66 x 51 x 12	67	30	DF4	DF4

*Indicates models that are MRI Conditional^{1,2}

Indications: The devices are intended to provide ventricular antitachycardia pacing and ventricular defibrillation for automated treatment of life-threatening ventricular arrhythmias. Contraindications: Contraindications for use of the pulse generator system include ventricular tachyarhythmias resulting from transient or correctable factors such as drug toxicity, electrolyte imbalance, or acute myocardial infarction.

Adverse Events: Implantation of the pulse generator system, like that of any other device, involves risks, some possibly life-threatening. These include but are not limited to the following: acute hemorrhage/bleeding, air emboli, arrhythmia acceleration, cardiac or venous perforation, cardiagenic shock, cyst formation, erosion, exacerbation of heart failure, extrusion, fibrotic tissue growth, fluid accumulation, hematoma formation, histotoxic reactions, infection, keloid formation, mocardial irritability, nerve damage, pneumothorax, thromboemboli, venous occlusion. Other possible adverse effects include mortality due to: component failure, device-programmer communication failure, lead abrasion, lead dislodgment or poor lead placement, lead fracture, inability to defibrillate, inhibited therapy for a ventricular tachycardia, interruption of function due to

electrical or magnetic interference, shunting of energy from defibrillation paddles, system failure due to ionising radiation. Other possible adverse effects include mortality due to inappropriate delivery of therapy caused by: multiple counting of cardiac events including T waves, P waves, or supplemental pacemaker stimuli. Among the psychological effects of device implantation are imagined pulsing, dependency, fear of inappropriate pulsing, and fear of losing pulse capability.

Refer to the User's Manual for detailed indications, contraindications, warnings, precautions and potential adverse events.

†QHR is a trademark of Greatbatch Medical





Compatible

Dovice Testing (Industion Methods

Ellipse[™] VR

Single-Chamber Implantable Cardioverter Defibrillator (ICD)

Product Specifications

PHYSICAL SPECIFICATIONS			Device Testing/Induction Methods	
Aodels Felemetry Delivered/Stored Energy (J)	CD1377-36C RF 36/39	CD1377-36QC RF 36/39	DC Fibber [™] Pulse Duration (sec) Burst Fibber Cycle Length (ms) Noninvasive Programmed	0,5-5,0 20-100
/olume (cc)	31	30	Stimulation (NIPS)	2-25 stimuli with up to 3 extrastimuli
Veight (g)	66	67	Patient Notifiers	
Size (mm)	68 x 51 x 12	66 x 51 x 12		
Defibrillation Lead Connections	DF1	DF4	Programmable Notifiers (On; Off)	Device at ERI; Charge Time Limit Reached; Possible HV Circuit Damage
Sense/Pace Lead Connections	IS-1	DF4		Ventricular Lead Impedance Out of Range; High-Voltage Lead Impedan Out of Range; %V pacing; CorVue [™] Congestion Trigger; SecureSense le
ligh-Voltage Can	Electrically active titanium can	Electrically active titanium can		noise detected, non-sustained lead noise detected, ST Episodes
Coating	Parylene No	Parylene		(Type I only)
/RI Conditional		Yes-MRI-ready	Device Parameter Reset	On
PARAMETER	SETTINGS		Entry into Backup VVI Mode	On
ensing/Detection			Vibration Duration (sec)	2; 4; 6; 8; 10; 12; 14; 16
Sense <i>Ability</i> ™ Technology	Automatic Sensitivity Control adjust	tment for ventricular events	Number of Vibrations per Notification Number of Notifications	1-16
ow Frequency Attenuation	On; Off		Time Between Notifications (hours)	10; 22
hreshold Start	(Post-Sensed; Ventricular) 50; 62,5			10, 22
	(Post-Paced; Ventricular) Auto; 0,2-		Electrograms and Diagnostics	
)ecay Delay	(Post-Sense/Post-Pace; Ventricular)) 0-220	Stored Electrograms	Up to 45 minutes including up to one minute programmable pre-trigger
/entricular Sense Refractory (ms) Detection Zones	125; 157 3 zone programming - 1 zone, 2 zone	an ar 2 Januar (VT 1 VT 2 VE)		data per VT/VF diagnosis/detection electrograms; triggers include:
SVT Discriminators	Sudden Onset, Interval Stability; Sir			diagnosis; detection; therapy; PC shock delivery; noise reversion;
SVI DISCITITIZATIS	Discrimination (Far Field MD or Orig			magnet reversion; and morphology template verification; lead noise
	or Automatic Template Update			detected, non-sustained lead noise detected, NSVT/NSVF
Discrimination modes	On, Passive, Off		Therapy Summary	Diagram of therapies delivered
SVT Threshold	150-240 min ⁻¹		Episodes Summary	Directory listing of up to 60 episodes with access to more details including stored electrograms
SVT Timeout	0,25-5 min		Lifetime Diagnostics	History of bradycardia events and device-initiated charging
Nonitor Mode	Detection, discrimination and diagn	ostics, no therapy delivery		
	(VT or VT-1 zone)		Histograms	Event Histogram; Ventricular Heart Rate Histogram; Exercise and
Reconfirmation	Continuous sensing during charging		mstograms	Activity Trending; DirectTrend™ reports up to 1 year
ead Noise Discrimination	SecureSense [™] RV lead noise discrin	nination	Real-Time Measurements (RTM)	Pacing lead impedances; high-voltage lead impedances;
	(On; On with Timeout; Passive; Off)		,	and signal amplitudes
ntitachycardia Pacing Therapy			ST Monitoring	ST Histogram Data; Long-term ST Deviation Trend; ST Episode Log;
TP Configurations	Ramp; Burst; Scan; 1 or 2 schemes			ST Episode Details; 24-Hour ST & HR Trend; ST EGM Baseline & Snapshot prior to ST Episode, VT/VF, Interrogation (Snapshots and 24-hour trend
TP in VF Zone	ATP While Charging; ATP Prior to Cha	arging; Off		at time of interrogation)
TP Upper Rate Cutoff	150 - 300 min ⁻¹		CorVue [™] Congestion Monitoring	On; Off
Burst Cycle Length Ain. Burst Cycle Length (ms)	Adaptive; Readaptive or Fixed 150-400 in increments of 5		CorVue Congestion Trigger	8-18 days
lumber of Bursts	1-15			
lumber of Stimuli	2-20			
ldd Stimuli per Burst	On: Off			
TP Pulse Amplitude (V)	7,5 Independent from Bradycardia a	nd Post-Therapy Pacing		
TP Pulse Width (ms)	1,0 or 1,5 Independently programma			
	and Post-Therapy Pacing			
ligh-Voltage Therapy				
)ynamicTx™ Algorithm	On; Off			
0eFT Response™ Technology	Programmable pulse width for P1/P2	2 and tilt		
ligh-Voltage Output Mode	Fixed Pulse Width; Fixed Tilt			
Vaveform	Biphasic; Monophasic			
RV Polarity	Cathode (-); Anode (+)	2		
lectrode Configuration	RV to Can; RV to SVC/Can; RV to SV	L .		
Bradycardia Pacing	0// 10////D			
Permanent Modes	Off; VVI(R)			
emporary Modes	Off; VVI; VOO			
Rate-Adaptive Sensor Programmable Rate Parameters	On, Off, Passive Off; Base Rate (min-1); Rest Rate (min	n-1), Maximum Sonsor Pato (min-1)		
rogrammable Rate Parameters		th (RV) (ms); Hysteresis Rate (min ⁻¹);		
	Rate Hysteresis with Search	th (hv) (his), hysteresis hate (hill),		
/entricular AutoCapture™	Nate Hysteresis with ocuron			
Pacing System	On; Off			
Post-Therapy Pacing (Independent	ly programmable from Bradycardia	and ATP)		
	Off; VVI			
Post-Shock Pacing Mode				
Post-Shock Pacing Mode Post-Shock Base Rate (min ⁻¹)	30-100 in increments of 5			

1. MRI Conditional Parameters: 1,5 Tesla, 2 W/Kg SAR

2. See MRI Procedure Information for approved MR Conditional Systems Device/Lead combinations and scan parameters

Customer Support: 46-8-474-4756

Brief Summary: Prior to using these devices, please review the Instructions for Use for a complete listing of indications, contraindications, warnings, precautions, potential adverse events and directions for use. Devices depicted may not be available in all countries. Check with your St. Jude Medical representative for product availability in your country.

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Durata[™] Defibrillation Lead

Product Highlights

- Allows patients to safely undergo an MRI scan when used in combination with an SJM MRI Ready device.^{1,2}
- Optim[™] insulation is a chemical co-polymer that offers superior handling and durability³
- Two innovative designs are intended to help prevent tissue ingrowth flatwire technology provides a low profile for the defibrillation coils, and silicone backfilling completely fills the shock coil space
- Redundant conductors serve as a backup system in the unlikely event of a conductor failure
- Symmetrically aligned cables within the lead body and centrally located coil provide for additional protection to the inner coil⁴
- The DF4 connector is designed to simplify implants by streamlining defibrillation connections into a single terminal pin and reducing the number of set screws

Ordering Information

Contents: Defibrillation lead

Model Number	Insulation	Fixation	Min. Introducer (F)	Shock Configuration	Sensing	Tip-to-Proximal Coil (cm)	Connector	Lengths (cm)
7120	Optim	Ext/Ret Helix	7	Dual-coil	True bipolar	17	DF1; IS-1	60; 65
7120Q	Optim	Ext/Ret Helix	7	Dual-coil	True bipolar	17	DF4	52; 58;*65*
7121	Optim	Ext/Ret Helix	7	Dual-coil	True bipolar	21	DF1; IS-1	60; 65; 75
7121Q	Optim	Ext/Ret Helix	7	Dual-coil	True bipolar	21	DF4	52; 58; 65
7122	Optim	Ext/Ret Helix	7	Single-coil	True bipolar	N/A	DF1; IS-1	60; 65; 75
7122Q	Optim	Ext/Ret Helix	7	Single-coil	True bipolar	N/A	DF4	52; 58;*65*
7170	Optim	Tines	7	Dual-coil	True bipolar	17	DF1; IS-1	60; 65; 75
7170Q	Optim	Tines	7	Dual-coil	True bipolar	17	DF4	52; 58; 65
7171	Optim	Tines	7	Dual-coil	True bipolar	21	DF1; IS-1	60; 65; 75
7171Q	Optim	Tines	7	Dual-coil	True bipolar	21	DF4	52; 58; 65
7172Q	Optim	Tines	7	Single-coil	True bipolar	N/A	DF4	52; 58; 65

*Indicates models and lead lengths that are MRI Conditional^{1,2}

Indications for Use: The Durata[™] transvenous leads are indicated for use with compatible pulse generators (refer to the applicable defibrillator manual for system indications). They provide pacing and sensing and deliver cardioversion/defibrillation therapy to the heart. A transvenous lead system may offer the patient the benefit of avoiding a thoracotomy for lead implantation. If the initial lead configuration is not effective, repositioning of the lead or other lead configuration should be attempted. In some patients, a nonthoracotomy lead configuration may not provide related conversion of arrhythmias, and the use of subcutaneous or epicardial patch defibrillation leads should be considered.

Contraindications: Contraindications for use of the Durata leads with an implantable pulse generator include ventricular tachyarrhythmias resulting from transient or reversible factors such as drug toxicity, electrolyte imbalance, or acute myocardial infarction. Transvenous lead systems are contraindicated for patients with tricuspid valual disease or a mechanical heart valve. Durata leads are contraindicated for patients for whom a single dose of 1.0 mg of dexamethasone sodium phosphate is contraindicated. The Durata leads are contraindicated for extra firm (red color knob) stylets. The lead is not designed, sold, or intended for use other than as indicated. St. Jude Medical DF1 lead connectors conform to the international connector standard ISO 11318/Amd.
St. Jude Medical IS-1 lead connectors conform to the international connector standard ISO 5841.
St. Jude Medical DF4 lead connectors conform to the international connector standard ISO 27186: 2010 (E).

Potential Complications: Possible complications of the use of transvenous lead systems include, but are not limited to, supraventricular or ventricular arrhythmias, conduction disturbances, cardiac perforation, cardiac tamponade, loss of contractility, air embolism, heart wall rupture, myocarditis, post-operative heart failure, chronic mechanical stimulation of the heart, tricuspid valve dysfunction, lead fracture necessitating surgical removal, pneumothorax, henothorax, infection, tissue necrosis and erosion of the skin. Specific events and effects are summarised below:

WARNING: Implanted cardiac leads are subjected to a hostile environment within the body due to constant, complex flexural and torsional forces, interactions with leads and/or the pulse generator, or other forces associated with cardiac contractions and patient physical activity, posture and anatomical influences Cardiac leads' functional lifetimes can be affected by these and other factors.

Refer to the defibrillator manual for additional complications and precautions specific to the pulse generator.





Durata™ **Defibrillation Lead**

Product Specifications PHYSICAL SPECIFICATIONS

True Bipolar, Active-Fixation Defibrillation Leads

Models	7120	71200	7121	7121Q	7122	71220
Fixation	Ext/Ret Helix	Ext/Ret Helix	Ext/Ret Helix	Ext/Ret Helix	Ext/Ret Helix	Ext/Ret Helix
Shock Configuration	Dual-Coil	Dual-Coil	Dual-Coil	Dual-Coil	Single-Coil	Single-Coil
Sensing Configuration	True Bipolar	True Bipolar	True Bipolar	True Bipolar	True Bipolar	True Bipolar
Min. Size Introducer	7 F	7 F	7 F	7 F	7 F	7 F
Lengths (cm)	60; 65	52; 58; 65	60; 65; 75	52; 58; 65	60; 65; 75	52; 58; 65
Connector	DF1; IS-1	DF4	DF1; IS-1	DF4	DF1; IS-1	DF4
Body Diameter	6,8 F	6,8 F	6,8 F	6,8 F	6,8 F	6,8 F
Tip-to-Anode Spacing	11 mm	11 mm	11 mm	11 mm	11 mm	11 mm
Tip-to-Proximal Coil	17 cm	17 cm	21 cm	21 cm	N/A	N/A
Tip Electrode Area	6 mm ²	6 mm ²	6 mm ²	6 mm ²	6 mm ²	6 mm ²
Steroid Plug	Yes	Yes	Yes	Yes	Yes	Yes
Distal Shock Coil Area	367 mm ²	367 mm ²	367 mm ²	367 mm ²	367 mm ²	367 mm ²
Proximal Shock Coil Area	588 mm ²	588 mm ²	588 mm ²	588 mm ²	N/A	N/A
MRI Conditional	No	Yes, MRI-ready (lengths: 58 and 65 cm)	No	No	No	Yes, MRI-ready (lengths: 58 and 65 cm

True Bipolar, Passive-Fixation Defibrillation Leads

Models	7170	7170Q	7171	7171Q	7172Q	
Fixation	Tines	Tines	Tines	Tines	Tines	
Shock Configuration	Dual-Coil	Dual-Coil	Dual-Coil	Dual-Coil	Single-Coil	
Sensing Configuration	True Bipolar					
Min. Size Introducer	7 F	7 F	7 F	7 F	7 F	
Lengths (cm)	60; 65; 75	52; 58; 65	60; 65; 75	52; 58; 65	52; 58; 65	
Connector	DF1; IS-1	DF4	DF1; IS-1	DF4	DF4	
Body Diameter	6,8 F					
Tip-to-Anode Spacing	11 mm					
Tip-to-Proximal Coil	17 cm	17 cm	21 cm	21 cm	N/A	
Tip Electrode Area	3.5 mm ²					
Steroid Plug	Yes	Yes	Yes	Yes	Yes	
Distal Shock Coil Area	367 mm ²					
Proximal Shock Coil Area	588 mm ²	588 mm ²	588 mm ²	588 mm ²	N/A	
MRI Conditional	No	No	No	No	No	

MRI Conditional Parameters: 1,5 Tesla, 2 W/Kg SAR
See MRI Procedure Information for approved MR Conditional Systems Device/Lead combinations and scan parameters
Jenney C, Tan J, Karicherla A, Burke J, Helland J. A New Insulation Material for Cardiac Leads with Potential for Improved Performance, Heart Rhythm, 2, S318-S319 (2005).
St. Jude Medical Engineering Report: Tension and Cable Shortening Comparison. Report 60032635

Customer Support: 46-8-474-4756

Brief Summary: Prior to using these devices, please review the Instructions for Use for a complete listing of indications, contraindications, warnings, precautions, potential adverse events and directions for use. Devices depicted may not be available in all countries. Check with your St. Jude Medical representative for product availability in your country.

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EC Certificate

EC Design-Examination Certificate

Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD), Annex 2 (4) (Other devices than custom made or intended for clinical investigation)

No. 17 17 07 14607 216

Manufacturer:

St. Jude Medical Cardiac Rhythm Management Division 15900 Valley View Court Sylmar, CA 91342



EC-Representative: St. Jude Medical Coordination Center BVBA

USA

The Corporate Village Da Vincilaan 11 Box F1 1935 Zaventem BELGIUM

Product:

Implantable Cardioverter / Defibrillators

The Certification Body of TÜV SÜD Product Service GmbH declares that a design examination has been carried out on the respective devices in accordance with AIMDD Annex 2 (4). This design of the devices conforms to the requirements of this Directive. For marketing of these devices an additional Annex 2 certificate is mandatory. See also notes overleaf.

Report no .:

713106728

Valid from: Valid until: 2017-09-26

2017-09-28

1. Pumil

Date, 2017-09-25

Stefan Preiß



TÜV SÜD Product Service GmbH is Notified Body with identification no. 0123

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EC Certificate EC Design-Examination Certificate Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD), Annex 2 (4) (Other devices than custom made or intended for clinical investigation)

No. 17 17 07 14607 216

Model(s):	see attachment
Parameters:	Ј.
Facility(ies):	St. Jude Medical Cardiac Rhythm Management Division 15900 Valley View Court, Sylmar, CA 91342, USA
	St. Jude Medical Puerto Rico LLC Lot A Interior - #2 Rd Km. 67.5, Santana Industrial Park, Arecibo PR 00612, USA
	St. Jude Medical Operations (M) Sdn.Bhd. Plot 102, Lebuhraya Kampung Jawa, Bayan Lepas Industrial Zone, 11900 Penang, MALAYSIA
	St. Jude Medical Coordination Center BVBA European Distribution Center, BruCargo 831, 1931 BruCargo, BELGIUM
Design Facility(ies):	St. Jude Medical Cardiac Rhythm Management Division 15900 Valley View Court, Sylmar, CA 91342, USA

Page 2 of 2



Attachment for Certificate no I7 17 07 14607 216 dated 2017-09-25

Product: Implantable Cardioverter / Defibrillators				
Test Report No.:	71362982			
Model:		Model No:		
Fortify™ VR		CD1233-40,	CD1233-40Q	
Fortify™ DR		CD2233-40,	CD2233-40Q	
Unify™		CD3235-40,	CD3235-40Q	
Test Report No.:	71376924			
Unify Quadra™		CD3251-40,	CD3251-40Q	
Test Report No.:	713000600/7	713000540		
Ellipse™ VR		CD1275-36,	CD1275-36Q,	
Ellipse™ DR		CD2275-36,	CD2275-36Q,	
Test Report No.:	713015987_1			
Quadra Assura™		CD3367-40,	CD3367-40C	
Quadra Assura MP	ТМ	CD3371 -4 0,	CD3371-40C	
Unify Assura™		,	CD3361-40C CD3361-40QC	
Fortify Assura™ DR	R	CD2359-40,	CD2359-40C	
Fortify Assura™ VF	२	CD1359-40,	CD1359-40C	
Ellipse™ DR		CD2377-36,	CD2377-36C	
Ellipse™ VR		CD1377-36,	CD1377-36C	

Page 1 of 2



Attachment for Certificate no I7 17 07 14607 216 dated 2017-09-25

Test Report No.: 713015987_1 / 713057341

Model:	Model No:	Variants:
Ellipse™ VR	CD1377-36Q, CD1377-36QC	MR Conditional
Ellipse™ DR	CD2377-36Q, CD2377-36QC	MR Conditional

Test Report No.: 713015987_1 / 713060615

Fortify Assura™ VR	CD1359-40Q,	CD1359-40QC	MR Conditional
Fortify Assura [™] DR	CD2359-40Q,	CD2359-40QC	MR Conditional

Test Report No.: 713015987_1 / 713068024

Quadra Assura™	CD3367-40Q,	CD3367-40QC	MR Conditional
Quadra Assura MP™	CD3371-40Q,	CD3371-40QC	MR Conditional

Munich, MHS-CRT, 2017-09-25

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Stefan Preiß Certification Medical Technology



90264657 Rev F **Declaration of Conformity**

SJM Declaration of Conformity Implantable Cardioverter/Defibrillators

St. Jude Medical (SJM) hereby declares that the following SJM facilities and products conform to the applicable provisions of Annex 2 of the European Union's Active Implantable Medical Devices Directive, AIMDD, 90/385/EEC. All supporting documentation is retained under the premises of SJM. We declare no application has been lodged with any other notified body for the same products. This declaration is issued under the sole responsibility of the manufacturer. This declaration supersedes any declaration issued previously for the same product(s).

Manufacturer Address:	St. Jude Medical Cardiac Rhythm Management Division 15900 Valley View Court Sylmar, CA 91342, USA
European Representative:	St. Jude Medical Coordination Center BVBA The Corporate Village Da Vincilaan 11 Box F1 1935 Zaventem, Belgium
Product Type:	Implantable Cardioverter/Defibrillators
Product Name(s):	See Attachment
Model Number(s):	See Attachment
Classification:	AIMD
GMDN Code(s):	See Attachment
Original CE Mark Date:	See Attachment
Certificate No. and expiration date:	EC Certification No: I7 17 07 14607 216 Expiration Date: 2022-09-25
	FQA Certificate No: I1 16 12 14607 211

Expiration Date: 2021-07-25 ISO13485

Certificate No: Q1N 17 09 14607 217 Expiration Date: 2020-10-31

Signature:

Kathy Berg

Manager Regulatory Affairs

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90264657 Rev F **Declaration of Conformity**

SJM Declaration of Conformity Implantable Cardioverter/Defibrillators

Applicable Quality System Standards:	Fulfills the requirements of Annex 2 of the European Union's Active Medical Devices Directive, AIMDD, 90/385/EEC/corresponding national legislation
	Fulfills applicable requirements including CE marking and the Essential Requirements of AIMDD, 90/385/EEC/corresponding national legislation
Notified Body:	TÜV SÜD Product Service GmbH Zertifizierstelle Ridlerstraße 65, 80339, Münich, Germany
Notified Body Number:	0123
Manufacturing Facilities:	St. Jude Medical Cardiac Rhythm Management Division 15900 Valley View Court Sylmar, CA 91342, USA
	St. Jude Medical Puerto Rico LLC Lot A Interior - #2 Rd Km. 67.5, Santana Industrial Park, Arecibo PR 00162, USA
	St. Jude Medical Operations (M) Sdn.Bhd Plot 102, Lebuhraya Kampung Jawa, Bayan Lepas Industrial Zone, 11900 Penang, MALAYSIA

Signature:

Kathy Berg

Manager Regulatory Affairs

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10V2017 **Issue Date**

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SJM Declaration of Conformity Implantable Cardioverter/Defibrillators ATTACHMENT TO DECLARATION OF CONFORMITY

The following product(s) is/are approved under EC-Certificate number 17 17 07 14607 216.

Product Name	Model No.	GMDN Code	First Date of CE Marking
Fortify™ VR	CD1233-40, CD1233-40Q	35852	2010-1-29
Fortify™ DR	CD2233-40, CD2233-40Q	37265	2010-1-29
Unify™	CD3235-40, CD3235-40Q	47270	2010-1-29
Unify Quadra™	CD3251-40, CD3251-40Q	47270	2011-3-15
Ellipse™ VR	CD1275-36, CD1275-36Q	35852	2012-2-3
Ellipse™ DR	CD2275-36, CD2275-36Q	37265	2012-2-3
Quadra Assura™ IS-1/DF-1	CD3367-40, CD3367-40C	47270	2012-12-18
Quadra Assura MP™ IS-1/DF-1	CD3371-40, CD3371-40C	47270	2012-12-18
Unify Assura™ IS-1/DF-1	CD3361-40, CD3361-40C CD3361-40Q, CD3361-40QC	47270	2012-12-18
Fortify Assura™ DR IS-1/DF-1	CD2359-40, CD2359-40C	37265	2012-12-18
Fortify Assura™ VR IS-1/DF-1	CD1359-40, CD1359-40C	35852	2012-12-18
Ellipse™ DR IS-1/DF-1	CD2377-36, CD2377-36C	37265	2012-12-18
Ellipse™ VR IS-1/DF-1	CD1377-36, CD1377-36C	35852	2012-12-18
Ellipse™ VR DF-4	CD1377-36Q, CD1377-36QC MR Conditional	35852	2015-05-11
Ellipse™ DR DF-4	CD2377-36Q, CD2377-36QC MR Conditional	37265	2015-05-11
Fortify Assura™ VR DF-4	CD1359-40Q, CD1359-40QC MR Conditional	35852	2015-7-14
Fortify Assura™ DR DF-4	CD2359-40Q, CD2359-40QC MR Conditional	37265	2015-7-14
Quadra Assura™ DF4	CD3367-40Q, CD3367-40QC MR Conditional	47270	2015-10-13
Quadra Assura MP™ DF-4	CD3371-40Q, CD3371-40QC MR Conditional	47270	2015-10-13

Signature:

Kathy Berg

Manager Regulatory Affairs

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